illumina

Analytical performance of the AmpliSeq[™] for Illumina Focus Panel with FFPE samples

Accurate, sensitive, and precise variant detection in tumor samples.

Introduction

Next-generation sequencing (NGS) has transformed cancer research, enabling variant detection in numerous genes in a single assay. By focusing on genes related to a particular disease or pathway, targeted sequencing provides the advantage of higher depth of coverage. This increased analytical sensitivity may be particularly important for detection of low frequency variants in heterogenous tumor tissues.

The AmpliSeq for Illumina Focus Panel was developed for biomarker analysis in both DNA and RNA concurrently. The panel enables highly accurate mutation detection in 52 genes with known relevance to solid tumors, including lung, colon, breast, ovarian, melanoma, and prostate (Table 1). Researchers can use the panel to investigate single nucleotide variants (SNVs), insertion/deletions (indels), and copy number variants (CNVs) in DNA samples, or gene fusions in RNA samples. The AmpliSeq for Illumina Focus Panel allows rapid and accurate assessment of genomic variation for translational and clinical research studies (Figure 1).

This application note demonstrates the high analytical sensitivity and precision of the AmpliSeq for Illumina Focus Panel for variant detection with reference standards and formalin-fixed, paraffin-embedded (FFPE) tumor samples using the Eppendorf ep*Motion®* in an automated library preparation workflow.

Table 1: AmpliSeq for Illumina Focus Panel gene content							
	DNA		R	NA			
Hotspot genes		CNVs	Fusior	n drivers			
AKT1	JAK1	ALK	ABL1	PPARG			
ALK	JAK2	AR	ALK	RAF1			
AR	JAK3	BRAF	AKT3	RET			
BRAF	KIT	CCND1	AXL	ROS1			
CDK4	KRAS	CDK4	BRAF				
CTNNB1	MAP2K1	CDK6	EGFR				
DDR2	MAP2K2	EGFR	ERBB2				
EGFR	MET	ERBB2	ERG				
ERBB2	MTOR	FGFR1	ETV1				
ERBB3	NRAS	FGFR2	ETV4				
ERBB4	PDGFRA	FGFR3	ETV5				
ESR1	PIK3CA	FGFR4	FGFR1				
FGFR2	RAF1	KIT	FGFR2				
FGFR3	RET	KRAS	FGFR3				
GNA11	ROS1	MET	MET				
GNAQ	SMO	MYC	NTRK1				
HRAS		MYCN	NTRK2				
IDH1		PDGFRA	NTRK3				
IDH2		PIK3CA	PDGFRA				



Figure 1: Demonstrated AmpliSeq for Illumina Focus Panel workflow—The AmpliSeq for Illumina Focus Panel is part of an integrated, streamlined NGS workflow that is compatible with automated library preparation, sequencing, and data analysis. *Indicates time required for sequencing-ready DNA and RNA combined library preparation.

Methods

Samples

Various reference and tumor FFPE samples were procured for use in this study. Reference samples included Quantitative Multiplex Reference Standard genomic DNA (gDNA) (Horizon Discovery Group, Cat no. HD701), OncoSpan gDNA (Horizon Discovery Group, Cat no. HD827), ALK-RET-ROS RNA fusion (Horizon Discovery Group, Cat no. HD784), Structural Multiplex Reference Standard gDNA (Horizon Discovery Group, Cat no. HD753), NA12878 (Coriell Institute for Medical Research), NCI-H2228 (American Type Culture Collection (ATCC), Cat no. CRL-5935), HL-60 (ATCC, Cat no. CCL-240), and the Seraseq Fusion RNA Mix v3 (SeraCare, Cat no. 0710-0431). DNA and RNA extracts from FFPE samples of varying tissue type and quality were obtained from the Illumina Biological Specimen Inventory (BSI) Systems repository (Table 2).

Table 2: Normal and tumor FFPE samples

DN	A	RNA		
Sample source	No. of samples	Sample source	No. of samples	
Bone	1	Bladder	1	
Brain	1	Brain	3	
Breast	4	Breast	4	
Colon	7	Colon	3	
Esophagus	2	Lung	7	
Lung	15	Ovary	1	
Pancreas	2	Prostate	3	
Prostate	2	Skin	3	
Skin	1	Stomach	1	
Stomach	3	Thyroid	3	
Cell lines	8	Unknown	1	
Total	46	Total	30	

Automated library preparation

Libraries were prepared from 10 ng of input RNA or DNA (or otherwise specified amounts), in accordance with the AmpliSeq for Illumina Focus Panel User Guide. Library preparation in this application note followed an automated protocol using the Eppendorf epMotion 5073m NGS solution. This system is capable of supporting up to 32 samples in a single run (16 DNA and 16 RNA samples). Automating library prep reduces potential variation and human error from multiple, repeated pipetting steps and provides less on-bench time. Reagents from the AmpliSeq for Illumina Library PLUS (96 Reactions, Cat no. 20019102), AmpliSeq for Illumina CD Indexes Set A (Cat no. 20019105), and AmpliSeq for Illumina Library Equalizer (Cat no. 20019171) were loaded on the robot as recommended by the automated methods. Resulting libraries were normalized (manually or automated with the Equalizer Kit) and pooled before loading on to a flow cell for sequencing, as recommended by the MiSeq or MiniSeq System Denature and Dilute Libraries Guide (Document # 15039740 v03 and Document # 100000002697 v07, respectively).

Sequencing

Prepared libraries were sequenced at 2×151 bp on the MiSeq System (Cat no. SY-410-1003) using the MiSeq Reagent Kit v3 (600 cycles, Cat no. MS-102-3003). Libraries were also run on the MiniSeq System (Cat no. SY-420-1001) using the MiniSeq High Output Kit (300 cycles, Cat no. FC-420-1003).

Data analysis

Resulting data was analyzed with the DNA Amplicon App, RNA Amplicon App, or the DNA + RNA Amplicon App, all available in BaseSpace[™] Sequence Hub. The DNA + RNA Amplicon App provides integrated analysis of DNA and/or RNA libraries through the DNA Amplicon app for somatic and germline variant calling, OncoCNV caller app for CNV detection, the RNA Amplicon App for fusion calling, and Pindel for structural variant detection. Confirmation of variants was conducted using digital droplet PCR (ddPCR) assay (BioRad) or an orthogonal NGS assay covering similar targets.

Results

Performance with FFPE samples

To demonstrate the analytical performance of the AmpliSeq for Illumina Focus Panel with FFPE tumor samples, 68 total FFPE samples (DNA and RNA) of varying quality obtained from BSI Systems were evaluated. After library prep and sequencing on the MiSeq System, analysis resulted in high performing sequencing metrics (Table 3, Figure 2). The AmpliSeq for Illumina Focus Panel provided highly accurate variant calling in FFPE samples, indicated by predictive metrics such as positive predictive value (PPV), negative predictive value (NPV), recall, sensitivity, and others (Table 4).

Table 3: Sequencing metrics^a for FFPE samples

	DNA		RÌ	A	
Metric	Mean	StDev	Mean	StDev	
On-target	96.4%	4.8%	68%	8%	
Coverage uniformity	93.8%	8.9%	_	_	
Mean amplicon coverage ^b	3631	1371	2836	560	

a. Sequencing metrics are from a representative MiSeq run.

b. The thresholds for metrics for RNA samples were 40% for on-target percentage and 1779 for mean amplicon coverage.



Figure 2: FFPE DNA aligned reads by sample quality—FFPE DNA sequenced on the MiSeq and MiniSeq Systems resulted in high percentage of on-target reads (blue dots and green rings) regardless of sample quality, measured by Δ Cq (triangles).

Table 4: Predictive value with FFPE samples

	SNV	Indel	CNV	Fusion ^a
True positive (TP)	19	10	16	3
False positive (FP)	0	0	0	0
True negative (TN)	17	18	13	5
False negative (FN)	0	0	0	0
Total	36	28	29	8
Positive predictive value (PPV) = TP/(TP+FP)	100%	100%	100%	100%
Negative predictive value (NPV) = TN/(TN+FN)	100%	100%	100%	100%
Recall = TP/(TP+FN)	100%	100%	100%	100%
Accuracy = (TP+TN)/(TP+FP+FN+TN)	100%	100%	100%	100%
False positive rate (FPR) = FP/N	0%	0%	0%	0%
False negative rate (FNR) = FN/(TP+FN)	0%	0%	0%	0%
Sensitivity = TP/(TP+FN)	100%	100%	100%	100%
Specificity = TN/(FP+TN)	100%	100%	100%	100%

a. FFPE RNA samples above alignment threshold were analyzed for fusion variants. Results showed confirmed detection of NTRK1-TPM3, EML4-ALK, and TMPRSS2-ERG in the qualified samples. Samples absent for EML4-ALK fusion were confirmed negative.

Variant calling accuracy and precision

To demonstrate the high accuracy of variant detection in DNA samples, references HD701 and HD827 were evaluated with the AmpliSeq for Illumina Focus Panel. Results show significant correlation between observed and expected variants above 5% variant allele frequency (VAF) threshold (official AmpliSeq for Illumina VAF cut-off), with 100% of variants (SNV and indels) detected for both standards (Figure 3).

To demonstrate the precision of the AmpliSeq for Illumina Focus Panel, intra-assay variability (correlation between replicates for the same user) and inter-assay variability (correlation between users) were determined using reference variants in HD701 and HD827 samples (Table 5). Repeatability and reproducibility ranged from R² correlation coefficient of 0.98 to 1.00 value.

HD/01 r	eterence	sample					
	User 1	User 2	User 3		Run 1	Run 2	Run 3
User 1				Run 1			
User 2	0.99			Run 2	0.98		
User 3	0.99	1.00		Run 3	0.99	0.99	
HD827 r	eference	sample					
	User 1	User 2	User 3		Run 1	Run 2	Run 3
User 1				Run 1			
User 2	1.00			Run 2	0.99		
User 3	0.99	1.00		Run 3	0.99	0.99	

Table 5	: Reproduci	bility and	repeatability for	reference	samples
110000					

Values represent correlation coefficient (R²) of comparison groups.

Gene fusion detection

To demonstrate the ability of the AmpliSeq for Illumina Focus Panel to recognize structural variants within RNA transcripts, reference samples HD784, NCI-H2228, and the Seraseq Fusion RNA Mix v3 were evaluated. Results showed a 100% call rate for gene fusions within these samples (data not shown).



Figure 3: Somatic variant calling above 5% threshold in reference samples—High correlation ($R^2 > 0.99$) was seen between expected and observed VAF for reference samples (n=20 per data point) with standard deviation.

Assay sensitivity

To demonstrate analytical sensitivity of the AmpliSeq for Illumina Focus Panel, a limit of detection (LOD) experiment was conducted with reference sample HD753. The amount of input gDNA used for library prep was titrated down from 20 ng to 1 ng. Resulting libraries were sequenced and assayed for variant calling across four replicates of

Prepare Library | Sequence | Analyze Data

753

				20 ng	10 ng	5 ng⁵	1 ng ^b	
Variant type ^a	Chromosome no.	Gene	Variant	% detected (replicates)				
SNV high GC	chr. 19	GNA11	Q209L	100% (4/4)	100% (4/4)	100% (4/4)	100% (4/4)	
SNV high GC	chr. 14	AKT1	E17K	100% (4/4)	100% (4/4)	100% (4/4)	100% (4/4)	
SNV low GC	chr. 3	PIK3CA	E545K	100% (4/4)	100% (4/4)	100% (4/4)	100% (4/4)	
Long insertion	chr. 7	EGFR	V769_D770ins ASV	100% (4/4)	100% (4/4)	100% (4/4)	100% (4/4)	
Long deletion	chr. 7	EGFR	∆E746–A750	100% (4/4)	100% (4/4)	100% (4/4)	100% (4/4)	
CNV	chr. 7	MET	amplification	100% (4/4)	100% (4/4)	75% (3/4)	0% (0/4)	
CNV	chr. 2	MYC-N	amplification	100% (4/4)	100% (4/4)	100% (4/4)	100% (4/4)	
a Variant informatio	on was provided by Horizon Di	scovery Group as p	art of the known genotype	of this reference sample	2			

a. Variant information was provided by Horizon Discovery Group as part of the known genotype of this reference sam
b. The AmpliSeq for Illumina Focus Panel assay recommends 10 ng sample input.

each input amount. With the exception of a CNV amplification in the *MET* gene, all variants assayed were called for all replicates across all dilutions (Table 7).

To further demonstrate assay sensitivity, titration experiments were performed in which reference samples with known mutations were titrated into normal (wildtype) reference samples. In the first experiment, gDNA from HL-60, a cell line with an amplification CNV resulting in 20 copies of the *MYC* gene, was titrated into a diploid background (NA12878 reference genome) to create DNA mixtures containing 20, 10, 8, 6, 4 and 2 copies (baseline) of the *MYC* gene. 10 ng of gDNA of each dilution were prepared, sequenced, and analyzed with the OncoCNV caller app. The AmpliSeq for Illumina Focus Panel enabled detection of all copies of the *MYC* gene, with significant correlation to CNV detection via ddPCR (Figure 4).





Two gene fusion LOD experiments were performed. In the first, the amount of input RNA extracted from NCI-H2228, positive for an EML4-ALK fusion, was titrated down from 10 ng to 0.05 ng input into a fusion negative sample (n=4 for each input). After library preparation, sequencing on the MiSeq System, and analysis with the RNA Amplicon app, gene fusion detection was evaluated (Table 8). In the second experiment, NCI-H2228 RNA was heat-degraded to assess fusion detection relative to sample quality as measured by DV150. A known fusion transcript was detected with high sensitivity at 20 copies, which was dependent on read alignment at \geq 40% (Table 9).

Table 8: Effect of input on EML4-ALK fusion detection

Input ^a (ng)	ddPCR copies	No. of expected fusion transcripts	% samples with ≥ 3 called	% samples with ≥ 2 called	% samples with ≥ 1 called
10	5440	3	100%	100%	100%
5	2660	3	100%	100%	100%
0.5	322	3	100%	100%	100%
0.25	130	3	100%	100%	100%
0.1	40	3	0%	50%	100%
0.05	20	3	0%	0%	100%
0	0	0	0%	0%	0%
a. Librar	ries were prep	bared using 10 ng t	otal RNA input wh	en combined wit	h a fusion

negative sample.

Table 9: Effect of sample quality on EML4-ALK fusion detection

DV150 ^a	No. of observed fusion transcripts (mean)	Aligned reads	Mean coverage	ddPCR copies
84	3	73%	4377	664
65	3	73%	2478	192
39	2	56%	3166	36
27	1	40%	1779	14.6
16	0	19%	892	2.6
9	0	1%	57	0
9	0.5	0	1	0

a. DV150 is the percentage of RNA fragments that are > 150 nucleotides in size.

Summary

The AmpliSeq for Illumina Focus Panel was developed for biomarker analysis in both DNA and RNA concurrently. The panel enables highly accurate mutation detection in 52 genes with known relevance to solid tumors. This application note demonstrates the high analytical sensitivity and precision of the AmpliSeq for Illumina Focus Panel for FFPE tumor sample variant detection using an automatedcompatible workflow.

Learn more

To learn more about the AmpliSeq for Illumina Focus Panel, visit www.illumina.com/products/by-type/sequencing-kits/library-prep-kits/ ampliseq-focus-panel.html.

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